JC02 Rec'd PCT/PTO 0 2 APR 2002

`					JUUZ Rec'd PCT/PTO 0 2 APR 2002			
Γ		PTCs 11-200		F COMMERCE PATENT AND TRADEMARK OFFICE	ATTORNEY'S DOCKET NUMBER 159-72			
r	TRANSMITTAL LETTER TO THE UNITED STATES U.S. APPLICATION NO. (If known, see 37 C F.R. 1.5)							
l				CTED OFFICE (DO/EO/US) ING UNDER 35 U.S.C. 371	10/ጫ&\$563			
ŀ	INTE	RNAT	IONAL APPLICATION NO.	INTERNATIONAL FILING DATE	PRIORITY DATE CLAIMED			
L			PCT/JP00/05170	02/08/2000	02/08/2000			
	TITL	E OF	INVENTION KITS FOR EXTRACING N	JCLEIC ACID AND METHOD OF EXTRAC	TING NUCLEIC ACID BY USING THE KITS			
ſ	APPLICANT(S) FOR DO/EO/US YOSHIHARA, N. et al.							
ľ	Appl	icant	herewith submits to the Unite	ed States Designated/Elected Office (DO/EC	0/US) the following items and other information:			
l	1.	Ø		of items concerning a filing under 35 U.S.C.				
l	2.	Ω.	This is a SECOND or SUBS	SEQUENT submission of items concerning a	filing under 35 U.S.C. 371.			
	3/	6		o begin national examination procedures (3	5 U.S.C. 371(f)). The submission must include			
l	4.	$\overline{\Box}$		by the expiration of 19 months from the prior	rity date (Article 31)			
١	5.	_		ation as filed (35 U.S.C. 371(c)(2)).	nty date (Article 01).			
l	Э.	a.	• •	uired only if not communicated by the Interr	national Bureau)			
l		a. b.		ed by the International Bureau.	ialional bureau).			
l				application was filed in the United States Re	people in a Office (PO/LIC)			
ļ	^	c. ⊠		application was lifed in the Office States reation of the International Application as filed				
١	6.	⊠	is attached hereto.	ation of the international Application as filed	(33 0.3.0. 37 1(c)(2)).			
l		a		ubmitted under 25 H S C 154(d)(4)				
ļ	_	b.		ubmitted under 35 U.S.C. 154(d)(4).	diale 10 (25 H S C 271/e)/2)\			
İ	7.	Π.		of the International Application under PCT A				
l		a.	_ ,	equired only if not communicated by the Inte	malional Bureau).			
١		b.	_	ated by the International Bureau.	andreamte has NOT ownized			
		C.		however, the time limit for making such ame	maments has NOT expired.			
١	_	d.	have not been made a		POT A#:-I- 10 (05 II C O 071(a)(0))			
l	8.			ation of the amendments to the claims under	PCT Article 19 (35 U.S.C. 371(c)(3)).			
l	9.	\boxtimes		e inventor(s) (35 U.S.C. 371(c)(4)).				
ļ	10.	П	A English language translat Article 36 (35 U.S.C. 3	ion of the annexes of the International Prelin 371(c)(5)).	minary Examination Report under PC1			
Ì	-4	Item	s 11 To 20 below concern	document(s) or information included:				
l	11.		An Information Disclosure S	Statement under 37 C.F.R. 1.97 and 1.98.				
l	12.	\boxtimes	An assignment document for	or recording. A separate cover sheet in com	pliance with 37 C.F.R. 3.28 and 3.31 is included.			
١	13.	\boxtimes	A FIRST preliminary amend	lment.				
Ì	14.		A SECOND or SUBSEQUE	NT preliminary amendment.	*			
1	15.		A substitute specification.	•				
١	16.		A change of power of attorr	ney and/or address letter.				
١	17.		A computer-readable form	of the sequence listing in accordance with P	CT Rule 13ter.2 and 35 U.S.C. 1.821-1.825.			
I	18.	$\overline{\Box}$		olished international application under 3				
I	19.	П	., .	sh language translation of the international a				
1	20.	⊠		PTO Form 1449, 3 sheets of drawings, and				
ı	20.	K.A.	Carro monito or anomiation.		- F-g Squonoo Elemiy			

-1-

JC13 Rec'd PCT/PTO 0 2 APR 2002

•										
U.S. APPLICATION NO. (II kno	wn, see 37 C.F.F.	1.5)	INTERNATIONAL AF		ON NO.		ATTC	RNEY'S DOCKET 159-72	NUME	BER
unkyow	4 7 7 1 4) /	PCT/JP00	J/U51/0			T C	ALCULATIONS	PTO	USE ONLY
21. The foftowing feit	ës are submit		(4) (E).				14	LULATIONS		
BASIC NATIONAL F	EE (37 C.F.H	. 1.49∠(a)((1)-(5): tion fee (37 C.F.R. 1.4	482)			ı			
			145(a)(2)) paid to USI				1			
nor international s	Search Reno	rt not pren:	ared by the EPO or Ji	PO		\$1040.00	l			
International preli	minarv exami	nation fee	(37 C.F.R. 1.482) not	t paid to			1			
USPTO but Intern	ational Searc	h Report p	repared by the EPO	or JPO.		\$890.00				
International preli	minary exami earch fee (37	nation fee C.F.R. 1.4	(37 C.F.R. 1.482) not 445(a)(2)) paid to USI	t paid to PTO		\$740.00				
International preli	minary exami	nation fee	(37 C.F.R. 1.482) pai PCT Article 33(1)-(4)	id to US	PTO					
						\$710.00	ı			
and all claims sat	mınary exami isfied provisio	nation fee ins of PCT	(37 C.F.R. 1.482) pai Article 33(1)-(4)	to US		\$100.00	L			
v	•		ENTER APPROP	RIATE	BASIC FE	E AMOUNT =	\$	890.00		
Surcharge of \$130.00 fo	r furnishing th	e oath or o	declaration later than	□ 20	□ 30		H		<u> </u>	
months from the earliest	claimed prior	ity date (3	7 C.F.R. 1.492(e)).				\$	0.00	L	
CLAIMS	NUMBER		NUMBER EXT	RA		ATE	1	0.00		
Total Claims	14	-20 =		-	X	\$18.00	\$	0.00	├	
Independent Claims	2	-3 =				\$84.00 80.00	\$	0.00	\vdash	
MULTIPLE DEPENDEN CLAIM FEES ARE NOT	I CLAIMS(S)	(II applica	DIE)	E ABO		LATIONS =	\$	890.00	-	
Applicant claims sr	nall ontity etc	HIS SAC 2	7 CFR 1.27. The fee			LA HONS =	┼╸	390.00	_	
are reduced by 1/2		us. 3663	, 5, 11 1.27. 1110 100	, maica	45046		1	0.00	l	
are reduced by 1/2						SUBTOTAL =	\$	890.00		
Processing fee of \$130.0	00, for furnish	ing the En	glish Translation later	r than			T	0.00		
months from the earliest	claimed prior	ity date (3	/ C.F.R. 1.492(t)).	T/	TAL NAT	ONAL FEE =	\$	890.00	-	
Fee for recording the en	closed assign	ment (37 (CFR 1 21(h)) The				+*	030.00	 	
accompanied by an app	ropriate cover	sheet (37	C.F.R. 3.28, 3.31).	\$40.00 r	er property	· +	\$	40.00		
Fee for Petition to Reviv	e Unintention	ally Aband	loned Application (\$12	280.00 -	 Small Ent 	ity = \$640.00	\$	0.00		
	TOTAL FEES ENCLOSED =					\Box				
			Ā	mount to be:						
							+	refunded	\$	
							┖	Charged	\$	
b. Please charge A duplicate cc The Commiss overpayment d. The entire cor application.	b. Please charge my Deposit Account No. 14-1140 in the amount of \$ to cover the above fees. A duplicate copy of this form is enclosed. The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 14-1140. A duplicate copy of this form is enclosed. The entire content of the foreign application(s), referred to in this application is/are hereby incorporated by reference in this									
NOTE: Where an appr or (b)) must be filed an	opriate time nd granted to	limit unde restore ti	er 37 C.F.R. 1.494 or ne application to per	1.495 h nding s	as not bee tatus.	en met, a peti	tion	to revive (37 (C.F.R	. 1.137(a)
					0	1.	L	У		
SEND ALL CORRESPO	ONDENCE TO) :			4	X 000	·	' '		
LUNGINA MANDEDUNE					SIGNA	тинь				
NIXON & VANDERHYE										
Arlington, Virginia 2220										
Telephone: (703) 816-4					B. J. S	adoff				
					NAME					
					00.000			A	00	
					36,663	RATION NUME	SEP.	April 2, 20 Date	U 2	
					ncuis	TIATION NOW	LIT	Date		

13 AUG 2002 10/089563

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Patent Application of

YOSHIHARA, N. et al.

Atty. Ref.: 159-72

Serial No. 10/089,563

Group:

US National Phase of PCT/JP00/05170

Filed: April 2, 2002

Examiner:

For: KITS FOR EXTRACTING NUCLEIC ACID AND METHOD OF EXTRACTING

NUCLEIC ACID BY USING THE KITS

August 13, 2002

Assistant Commissioner for Patents Washington, DC 20231

Sir:

AMENDMENT

Responsive to the Notification dated June 13, 2002 (copy attached), entry and consideration of the following amendments and remarks are requested.

IN THE SPECIFICATION

Amend the specification as follows:

Insert the attached Sequence Listing in place of the originally-filed copy of the same

REMARKS

The specification has been amended to include the attached Sequence Listing.

YOSHIHARA, N. et al. Serial No. **10/089,563** US National Phase of PCT/JP00/05170

The attached paper and computer readable copies of the Sequence Listing are the same. No new matter has been added. A separate Statement to this effect is attached.

An early and favorable Action on the merits is requested.

Respectfully submitted,

NIXON & VANDERHYE P.C.

By:

B. J. Sadoff Reg. No. 36.663

1100 North Glebe Road, 8th Floor Arlington, VA 22201-4714 Telephone: (703) 816-4000 Facsimile: (703) 816-4100

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Patent Application of

YOSHIHARA, N. et al.

Attv. Ref.: 159-72

Serial No. unknown

Group:

US National Phase of PCT/JP00/05170

Filed: April 2, 2002

Examiner:

For: KITS FOR EXTRACTING NUCLEIC ACID AND METHOD OF EXTRACTING NUCLEIC ACID BY USING THE KITS

April 2, 2002

Assistant Commissioner for Patents Washington, DC 20231

Sir:

PRELIMINARY AMENDMENT

In order to place the above-identified application in better condition for examination, please amend the application as follows:

IN THE SPECIFICATION

Please substitute the following paragraphs in the specification for corresponding paragraphs previously presented. A copy of the amended specification paragraphs showing current revisions is attached.

Page 1, before the first line, please insert as a separate paragraph:

This application is the US national phase of international application PCT/JP00/05170 filed 02 August 2000, which designated the US.

YOSHIHARA, N. et al. Serial No. **unknown** US National Phase of PCT/JP00/05170

IN THE CLAIMS

Please substitute the following amended claims for corresponding claims previously presented. A copy of the amended claims showing current revisions is attached.

- (Amended) The nucleic acid isolation kit of Claim 1 wherein the nucleic acid is RNA.
- (Amended) The nucleic acid isolation kit of Claim 1 wherein the reducing agent is 2-mercaptoethanol or dithiothreitol.
- (Amended) The nucleic acid isolation kit of Claim 1 wherein the coprecipitant is glycogen or dextran.
- (Amended) The nucleic acid isolation kit of Claim 1 wherein the protein denaturant is guanidine thiocyanate.
- (Amended) The method of Claim 7 comprising adding a protein denaturant during alcohol precipitation.

YOSHIHARA, N. et al.

. Serial No. unknown

US National Phase of PCT/JP00/05170

- (Amended) The method of Claim 7 wherein the biological component is a body fluid or a blood product.
- 12. (Amended) The method of Claim 7 wherein the biological sample has a volume of 30 μ 1 to 100 μ 1.
 - 13. (Amended) The method of Claim 7 comprising no step of adding a salt.
- (Amended) The method of Claim 7 wherein said steps are performed in a single 0.5 ml tube.

YOSHIHARA, N. et al. 'Serial No. unknown US National Phase of PCT/JP00/05170

REMARKS

Attached hereto is a marked-up version of the changes made to the specification and claims by the current amendment. The attached page(s) is captioned "Version With Markings To Show Changes Made."

Respectfully submitted,

NIXON & VANDERHYE P.C.

B. J. Sadoff

Reg. No. 36,663

BJS:ecb 1100 North Glebe Road, 8th Floor Arlington, VA 22201-4714 Telephone: (703) 816-4000

YOSHIHARA, N. et al. Serial No. unknown

US National Phase of PCT/JP00/05170

VERSION WITH MARKINGS TO SHOW CHANGES MADE

IN THE SPECIFICATION

Page 1, before the first line, please insert as a separate paragraph:

This application is the US national phase of international application PCT/JP00/05170 filed 02 August 2000, which designated the US.

IN THE CLAIMS

- (Amended) The nucleic acid isolation kit of Claim 1-or 2 wherein the nucleic acid is RNA.
- (Amended) The nucleic acid isolation kit of any one of Claims 1-to-3
 wherein the reducing agent is 2-mercaptoethanol or dithiothreitol.
- (Amended) The nucleic acid isolation kit of any one of Claims 1 to 4
 wherein the coprecipitant is glycogen or dextran.
- (Amended) The nucleic acid isolation kit of any one of Claims 1+6-5
 wherein the protein denaturant is guanidine thiocyanate.
- (Amended) The method of any one of Claims 7 to 9-comprising adding a protein denaturant during alcohol precipitation.

YOSHIHARA, N. et al.
Serial No. unknown
US National Phase of PCT/JP00/05170

 (Amended) The method of any one of Claims 7-to 10 wherein the biological component is a body fluid or a blood product.

- 12. (Amended) The method of any one of Claims 7-to 10 wherein the biological sample has a volume of 30 μ l to 100 μ l.
- 13. (Amended) The method of any-one of Claims 7-to-11 comprising no step of adding a salt.
- 14. (Amended) The method of any one of Claims 7+o-12 wherein said steps are performed in a single 0.5 ml tube.

3/prts

SPECIFICATION

KITS FOR EXTRACTING NUCLEIC ACID AND METHOD OF EXTRACTING NUCLEIC ACID BY USING THE KITS

5 TECHNICAL FIELD

The present invention relates to nucleic acid isolation kits and nucleic acid isolation methods using said kits.

BACKGROUND ART

10

Detection of target nucleic acids such as specific DNA or RNA in biological samples such as blood and saliva is very important not only in the field of research but also in the clinical field.

For example, the rapid increase in the number of patients with human immunodeficiency virus (HIV) infection or AIDS in recent years is a matter of concern not only in Japan but also worldwide. A recent epidemiological tendency of HIV infection is a marked increase in the proportions of vertical infection or infection at younger 20 ages. In the clinical field, there is an earnest demand for the development of an accurate and rapid genetic diagnosis. In Japan, the main infection route of HIV infection or AIDS especially in hemophiliacs is thought to be transfusion of blood including blood products. However, little detail has been elucidated for the infection route from blood products (Lin Oi Zhang, Peter Simmonds, Christopher A. Ludlam and Andrew J. Leigh Brown (1991). AIDS, pp. 675-681). It is also important to follow up the

12

progress of HIV decrease in patients undergoing AIDS therapy.

Currently, an indirect detection method based on HIV antigen-antibody reaction has been established and become a 5 major diagnostic means in laboratories. However, detection based on antigen-antibody reaction has been associated with the problem of a window period especially in screening. This is a time-lag between antigen infection and antigen or antibody production. For early clinical diagnosis of HIV 10 infection, it is important to accurately, directly and rapidly detect the presence of HIV RNA in blood as an alternative to the indirect method based on antigenantibody reaction. It is highly desirable to develop a new detection method that can be performed on smaller amounts 15 of samples with higher sensitivity and at reduced cost and with a shortened window period. It is necessary to diagnose HIV infection from very small amounts of samples especially in infection in children or vertical infection.

20 possible with the new introduction of molecular biological techniques such as polymerase chain reaction (PCR) or screening, for example, into clinical applications (Gerald Schochetman and John J. Sninsky (1991), "Direct Detection of Human Immunodeficiency Virus Infection Using the
25 Polymerase Chain Reaction" Springer-Verlag pp. 90-110; Janet S. Bootman, Pete A. Kitchin (1994), J. Virological Methods, pp. 1-8; Anne-Mieke Vandamme, Sonia Van Dooren, Wessel Kok, Patrick Goubau, Katrien Fransen, Tim Kievits,

Direct and rapid assay of HIV itself has become

Jean-Claude Schmit, Erik De Clercq, Jan Desmyter (1995), J.
Virological Methods pp. 121-132; E. Lyamuya, U. BredbergRaden, J. Albert, O. Grankvist, V. Msangi, C. Kagoma, F.
Mhalu, and G. Biberfield (1997), J. Clinical Microbiology,

pp. 278-280). Such nucleic acid detection systems
combining nucleic acid isolation with nucleic acid
amplification reactions such as PCR consist of three
distinct processes: i.e., nucleic acid isolation,
amplification and detection. Nucleic acid isolation is an
important process for subsequent amplification and
detection, and if a nucleic acid isolation technique can be
improved, the efficiency of amplification and detection
will increase.

Currently, several types of total RNA isolation

methods are known and a plurality of kits are commercially available (John M. Chirgwin, Alan E. Przybyla, Raymond J. MacDonald and William J. Rutter (1979), Biochemistry pp. 5294-5299; Osamu Yamada, Toshiya Matsumoto, Masahiro Nakashima, Shinobu Hagari, Toshio Kamahora, Hiroshi Ueyama, Yuichiro Kishi, Hidetoshi Uemura and Takashi Kurimura (1990), J. Virological Methods pp. 203-210).

However, these methods have disadvantages including the use of organic solvents such as phenol or chloroform to remove proteins and the necessity of changing tubes many 25 times during the isolation process. Further, they are not ideal for isolation from very small amounts of samples, especially blood products or blood samples from children. They also have the disadvantage that costs add up when a

number of samples are assayed in screening or the like. Thus, it has become increasingly important to develop a rapid and inexpensive HIV RNA detection system.

In recent years, several types of methods for

isolating a nucleic acid, especially RNA have come to be
known such as the guanidinium cyanate-phenol-chloroform
(AGPC) method (Piotr Chomczynski, Nicolette Sacchi (1987),
Analytical Biochemistry pp. 156-159) and methods using
CsCl, or a resin (Maniatis T. et al.: Molecular Cloning: A
laboratory manual, 2nd ed., Cold Spring Harbor 1989).

Japanese Patent Public Disclosure No. 236499/1995
incorporated herein as reference discloses an improved AGPC
method for isolating a viral nucleic acid, in which the
isolation process can be completed in a single tube without
using an organic solvent.

SUMMARY OF THE INVENTION

20

An object of the present invention is to provide a nucleic acid isolation kit comprising a reducing agent, a coprecipitant and a protein denaturant characterized in that it contains no protease.

The nucleic acid isolation kit of the present invention is preferably an RNA isolation kit.

Another object of the present invention is to provide a method for isolating a nucleic acid from a biological sample, comprising:

 i) incubating the biological sample with a reducing agent, a coprecipitant and a protein denaturant to degrade and denature proteins and other contaminants in the biological sample without using a protease, and

ii) directly performing alcohol precipitation with a lower alcohol.

BRIEF DESCRIPTION OF DRAWINGS

5

15

- FIG. 1 is a set of electrophoretic pattern of reaction products after nucleic acid isolation at varying glycogen concentrations and amplification.
 - FIG. 2 is a flowchart of a preferred embodiment of a nucleic acid isolation method of the present invention.
- FIG. 3 shows the sequences of nucleic acids isolated by a method of the present invention.

DETAILED DESCRIPTION OF THE INVENTION

Methods for nucleic acid detection consist of three main processes, i.e., nucleic acid isolation, amplification and detection. Improvements in the nucleic acid isolation process are also important for the subsequent amplification and detection processes.

Before the present invention was made, the isolation technique used in many nucleic acid detection methods was the AGPC method described above. The present inventors found a novel method for isolating a nucleic acid from a sample by using an improved AGPC method in combination with a coprecipitant, to accomplish the present invention. The details are described below.

25 Nucleic acid isolation kits

Specifically, the present invention provides a nucleic isolation kit comprising a reducing agent, a coprecipitant and a protein denaturant characterized in that it contains no protease.

A key feature of the present invention is that no protease is used. In conventional methods, non-specific proteases such as proteinase K, pronase and subtilisin are 5 first used to remove proteins from biological samples. In the present invention, nucleic acids can be efficiently isolated only with a coprecipitant and a protein denaturant without using such a protease.

i) Reducing agents

10

15

20

In the present invention, reducing agents include, but are not limited to, 2-mercaptoethanol, dithiothreitol, etc. These reducing agents are used for protection of sulfhydryl groups in proteins or for reductive cleavage of disulfide bonds.

Reducing agents can be preferably used at a concentration of, but not specifically limited to, about Reducing agents may be added to nucleic acids as premixes in protein denaturants.

ii) Coprecipitants

In the present invention, coprecipitants are not specifically limited to, and any coprecipitants that can function as carriers for coprecipitating nucleic acids are suitable. Preferred examples of coprecipitants are macromolecular polysaccharides such as glycogen and 25 dextran.

Glycogen is a polymer of D-glucose having a molecular weight of one million to several millions and is known as one of the most potent carriers for nucleic acid

precipitation (Steven Tracy (1981), Biochemistry pp. 251-268), and several types of commercially available glycogen can be used for in vitro applications. In the present invention, glycogen from slipper limpet is preferred from the viewpoints of cost and detection sensitivity.

The concentration of coprecipitants is not specifically limited, but as an example, glycogen from slipper limpet is used at 0.1 mg/ml to 2.0 mg/ml, preferably 0.2 mg/ml to 2.0 mg/ml, most preferably about 0 0.2 mg/ml (Example 2).

iii) Protein denaturants

20

2.5

Suitable protein denaturants are those known to be able to solubilize proteins. They include, but not specifically limited to, guanidine thiocyanate and urea.

15 Guanidine thiocyanate is especially preferred. A preferred embodiment of the present invention uses glycogen as a coprecipitant and guanidine thiocyanate as a protein denaturant, and is called the glycogen-guanidinium (GG) method.

Protein denaturants can be used at a final concentration of 3 M to 7 M, preferably 4 M to 6 M, most preferably approximately 4.5 M. Preferably, protein denaturants at the saturation concentration can solubilize proteins and efficiently reduce RNases that may be included in biological samples particularly in the case of isolation of RNA.

- 7 -

Preferably, protein denaturants are added to reactants after biological samples are mixed with coprecipitants. As described above, they may be added as premixes with reducing agents to reaction solutions.

Further, without ant limitation, protein denaturants can be optionally added to an alcohol during alcohol precipitation at the final stage of the isolation process to more clearly remove contaminant proteins especially in the case of samples containing a lot of contaminants such as blood products. Protein denaturants are added to an alcohol during the alcohol precipitation step preferably at 0.5 M to 2.0 M, most preferably about 0.9 M (Example 3).

In the present invention, there is no special need to externally add a salt during the nucleic acid isolation process, and nucleic acid isolation kits specially require no salt. Moreover, the pH need not be specially externally adjusted.

Nucleic acid isolation methods

20

The present invention also provides methods for isolating a nucleic acid from a biological sample.

Methods of the present invention comprises:

- incubating the biological sample with a reducing agent, a coprecipitant and a protein denaturant to degrade and denature proteins and other contaminants in the biological sample without using a protease, and
- 11) directly performing alcohol precipitation with a 25 lower alcohol.

In the present invention, nucleic acid isolation can be normally made from biological samples in a volume of about 50 μ l, preferably 30 μ l to 100 μ l. This means that

significantly smaller amounts of biological samples are required than conventional methods.

According to a preferred but non-limitative
embodiment, a biological sample is mixed with a

5 coprecipitant and then a reducing agent and a protein
denaturant are added in step i).

In step i), incubation takes place at 55°C - 65°C, preferably about 60°C for 5 minutes to 15 minutes, preferably about 10 minutes. Most preferably, incubation 10 takes place at about 60°C for about 10 minutes. Incubation can be performed in an apparatus such as, but not limited to, an incubator, PCR apparatus (eg, Thermal Cycler® from Perkin-Elmer), etc.

In step ii), the reaction mixture obtained in step i)

is then combined with a lower alcohol to precipitate a
nucleic acid. Alcohol precipitation can be performed by
known techniques. Suitable alcohols include isopropanol
and ethanol or the like. Preferably, isopropanol is added
at a final concentration of 40% or more or ethanol is added
at a final concentration of 70% or more. Then, the mixture
may be cooled at -70°C to 4°C after addition of a lower
alcohol to promote efficacy of salting out.

A protein denaturant may be added during alcohol precipitation as described above. This can remove contaminants precipitating with a target nucleic acid.

Alcohol precipitation is followed by centrifugation at 14,000 x g to 19,000 x g for 5 minutes to 15 minutes. Centrifugation may be performed at room temperature or

under cooling at about 4°C. After centrifugation, the supernatant is decanted or sucked to recover the precipitated nucleic acid.

Finally, the obtained nucleic acid can be washed with 5 70% ethanol, for example, and redissolved in a suitable solution into a ready-to-use state.

above can be performed in a single tube. This can prevent contamination which may otherwise be caused by a change of tubes during the isolation process. This is especially important particularly when the nucleic acid is amplified by PCR or the like. Moreover, the isolation process of the present invention can be performed in a 0.5 ml tube in contrast to conventional methods that required a tube of at least 1.5 ml or more in size. Thus, the necessary amount of each reagent can be reduced and also the nucleic acid redissolved in a 0.5 ml tube can be directly used for amplification reaction.

A preferred but non-limitative embodiment of the
present invention is shown in Example 1 and Fig. 2.

Nucleic acid isolation methods of the present invention described above can be applied to isolate both DNA and RNA. Preferably, they can be used for RNA isolation. In addition to RNA of the AIDS virus HIV described in the examples below, they can be clinically applied to type C hepatitis virus (HCV), influenza virus, type A hepatitis virus and the like. In addition to blood samples such as serum and plasma, they are also useful for

soluble liquid samples such as cerebrospinal fluid, saliva, semen and urine. They can also be used for powdered blood products derived from plasma or serum so that they have wide applications. Particularly, they are effective for small amounts of samples such as samples from children or powdered blood products.

Nucleic acids isolated by methods of the present invention can be subsequently amplified and detected, if necessary. That is, methods of the present invention can be combined with amplification techniques such as PCR to detect even very small amounts of nucleic acids at low concentrations that could not be detected by conventional methods.

Amplification

10

15

20

25

(1) Reverse transcription reaction

When the target nucleic acid is an RNA such as viral RNA, the RNA can be reversely transcribed with a reverse transcriptase capable of converting the RNA into cDNA directly in the tube used for isolation. That is, the RNA isolated by methods of the present invention is highly pure and contains no substances inhibiting reverse transcription reaction. Suitable reverse transcriptases include, for example, a reverse transcriptase derived from avian myeloblastosis virus used in Example 1 described below.

(2) Amplification of nucleic acids

Nucleic acids isolated by methods of the present invention can be amplified by known nucleic acid amplification reactions such as polymerase chain reaction (PCR) using primers specific to the nucleic acids directly in the case of DNA or after reverse transcription into cDNA in the case of RNA. Nucleic acids isolated by methods of the present invention are highly pure and they contain no substances inhibiting amplification reaction even though ordinary PCR techniques require strict conditions.

Amplification reaction of nucleic acids may be nested PCR involving first PCR with an outer primer pair of the target nucleic acid followed by second PCR with an inner primer pair as described in Example 1 below, for example. The extent of amplification may vary with the affinity of the primer for the target nucleic acid. In HIV-1 described in the examples below, for example, gag primers and pol primers gave somewhat different extents of amplification and pol primers showed a higher detection sensitivity than gag primers.

Detection

25

Amplified nucleic acids can be detected and identified by standard techniques such as electrophoresis and sequencing. For example, they can be detected by ethidium bromide staining after electrophoresis on a polyacrylamide gel or an agarose gel. The sequence can also be identified by known methods using commercially available nucleic acid sequencers.

As described above, methods of the present invention achieve high cost efficiency, high sensitivity, safety and time reduction in nucleic acid isolation. They can be combined with amplification techniques such as PCR to provide accurate, direct and rapid diagnosis for virus infection, for example. The GG method of the present invention can be applied not only to the field of research but also to the clinical field.

The following examples further illustrate the present invention but are not intended to limit the technical scope of the invention. Those skilled in the art can readily add modifications/changes to the present invention on the basis of the description of the specification, and those

10 modifications/changes are included in the technical scope of the present invention.

EXAMPLES

In the examples of the present invention, the following materials were used unless otherwise specified.

15 Materials

(1) Samples

In both AMPLICOR HIV-1 MONITOR® Test Kit (Roche) and NASBA Amplification System (Organon Teknika, Boxtel, Netherlands), 26 samples containing high copy numbers of HIV-1 RNA (30,000 copies or more / ml) and 47 samples containing low copy numbers below the detection limit (1,000 copies or less / ml) were used.

(2) Glycogen

Oyster glycogen (Sigma), slipper limpet glycogen

25 (Sigma), rabbit glycogen (Sigma) or bovine glycogen (Sigma)
was used.

(3) RNA isolation methods used as controls The following commercially available 7 kits/methods were used as controls of the present invention in the examples.

TRIZOL LS®

ISOGEN LS® (Nippon Gene)

5 RNA Isolation Technique (Stratagene)

SepaGene-RV® (Sanko Junyaku)

NASBA RNA Isolation Kit (Organon Teknika)

Smitest® (SUMITOMO METAL INDUSTRIES., LTD)

Cartrimox (Iowa Biotechnology).

10 Example 1: RNA isolation by the GG method and amplification and detection of the isolated RNA

HIV-1 RNA was isolated, amplified and detected from HIV-1-containing samples by the following procedures.

a. RNA isolation

15

RNA was isolated from samples following the scheme shown in Fig. 2 as an example of the GG method of the present invention.

In a 0.5 ml tube, 1 μ l of glycogen (10 mg/ml) was added to 50 μ l of each sample and the tube was agitated. 20 The tube was incubated at 60°C for 10 minutes with 150 μ l

- of 6 M guanidine thiocyanate containing 1% 2mercaptoethanol (2ME). Then, 200 µl of isopropanol was added. The tube was centrifuged at 15,000 g for 15 minutes at room temperature, and then the supernatant was removed.
- 25 400 µl of 70% ethanol containing 0.9 M guanidine thiocyanate was added. The tube was centrifuged at 15,000 g for 5 minutes at room temperature, and then the supernatant was removed. 400 µl of 70% ethanol was added.

The tube was centrifuged at 15,000 g for 5 minutes at room temperature, and then the supernatant was removed. The residue was dissolved in 10 µl of RNase-free sterile water.

b. Amplification

1) Primers

For amplification of HTV-1 RNA, gag and/or pol primers (SEQ ID NOS: 1-8) shown in Table 1 below were used (derived from a human retrovirus of AIDS obtained from Los Alamos National Laboratory).

Table 1

10

Primer Sequence Location SEO ID NO:
GF62 AAGGATAGAGGTAAAAGACACCA gag 270-292 1

gag 270-292 GF63 TAGCTGCTGGTCCAATGCTTTTA gag 1022-999 2 gag 581-599 15 SK100 ATCAAGCAGCCATGCAAAT gag 871-859 SK104 CTTTTGGTCCTTGTCTTATGTC pol 2385-2407 5 UNIPOL 1B AGTGGATATATAGAAGCAGAAGT UNIPOL 1A CCCCCAATCCCCCTTTTCTTTTAAAA pol 2722-2696 6 pol 2453-2470 7 P5 ATTAGCAGGAAGATGGCC pol 2594-2577 8 P6 TACTCCTTGACTTTGGGG 20

The primer pair GF62 and GF63 and the primer pair
UNIPOL 1B and UNIPOL 1A are outer primer pairs of gag and
pol, respectively. On the other hand, the primer pair
25 SK100 and SK104 and the primer pair P5 and P6 are inner
primer pairs of gag and pol, respectively.

2) PCR

The RNA isolated as above was used for nucleic acid

amplification by a conventional PCR method.

- i) Initially, 2 drops of mineral oil was added to a 0.5 ml tube containing the above isolated RNA dissolved in 10 μ l of sterile water and the tube was momentary spun.
- 5 Then, the tube was incubated at 80°C for 10 minutes in a Thermal Cycler® (Perkin-Elmer) to destroy any RNA complexes or secondary structures that may disturb priming in PCR.

 After 10 minutes, the tube was immediately transferred into ice to stop the reaction and momentary spun.
- 10 ii) Then, 15 µl of a mixture for RT having the composition shown in Table 2 below was added and the tube was vortexed and momentary spun.

Table 2. Composition of mixture for RT

15	H ₂ O	7.13
	x 10 PCR buffer	2.5
	10 mM MgCl ₂	2.0
	10 mM DTT	0.5
	2.5 mM dNTP	1.5
20	.1 μl/ml Primer 1 (outer)	0.25
	1 µ1/ml Primer 2 (outer)	0.25
	10 U/μl RNase inhibitor	0.77
	2.5 U/µl Reverse transferase (AMVRT)	0.1
	Total	15.00 µl

25

The tube was incubated in a warm bath (42°C) for 60 minutes to synthesize cDNA from mRNA. Then, the tube was momentary spun.

Þ

iii) Then, the tube was incubated at 99°C for 6 minutes in a Thermal Cycler to denature double stranded mRNA-cDNA into single strands. After 6 minutes, the tube was immediately transferred into ice to stop the reaction and momentary spun.

1v) Then, 75 $\mu \hat{L}$ of a mixture for first PCR having the composition shown in Table 3 below was added and the tube was vortexed and momentary spun.

10	Table	з.	Composition	of	mixture	for	first	PCR

H ₂ O	58
x 10 PCR buffer	7.5
10 mM MgCl ₂	3.0
2.5 mM dNTP	4.5
1 µl/ml Primer 1 (outer)	0.75
1 µl/ml Primer 2 (outer)	0.75
Taq polymerase	0.5
Total	75 00 u1

15

20

25

First PCR was performed in a Thermal Cycler. Specifically, 30 cycles of 94°C for 30 seconds, 55°C for 30 seconds and 72°C for 1 minute were performed. Then, the tube was incubated at 72°C for 10 minutes and then, the

reaction was stopped at 4°C.

v) Then, 10 μ l of the first PCR product obtained in iv) and mineral oil were added to a fresh 0.5 ml tube, and the tube was momentary spun. Then, the tube was incubated at 99°C for 6 minutes in a Thermal Cycler to denature

double-stranded DNA into single strands. After 6 minutes, the tube was immediately transferred into ice to stop the reaction and momentary spun.

vi) Then, 90 µl of a mixture for second PCR having the composition shown in Table 4 below was added and the tube was vortexed and momentary spun.

Table 4. Composition of mixture for second PCR

	H ₂ O	66.5
10	x 10 PCR buffer	10
	10 mM MgCl ₂	5
	2.5 mM dNTP	6
	1 μl/ml Primer 3 (inner)	1
	1 μl/ml Primer 4 (inner)	1
15	Taq polymerase	0.5
	Total	90.0 µl

Second PCR was performed in a Thermal Cycler.

Specifically, 30 or 40 cycles of 94°C for 30 seconds, 55°C

for 30 seconds and 72°C for 1 minute were performed. Then,
the tube was incubated at 72°C for 10 minutes and then, the
reaction was stopped at 4°C.

c. Detection

25

Electrophoresis and staining

Following amplification, PCR products were electrophoresed. Specifically, a dye was added to the second PCR product and an aliquot of 10 µl was loaded on a 5% acrylamide gel or a 2% agarose gel and electrophoresed.

HaeIII digest of ФX174 (Roche) was used as a molecular weight marker. Culture supernatant of an HIV-1 infected cell line MOLT-4/HTLV III was used as a positive control.

After completion of migration, bands were visualized by ethidium bromide staining.

Example 2: Optimal concentration of glycogen used for RNA isolation

The concentration of glycogen used as a carrier was diversified to determine the optimal concentration from the 10 viewpoints of isolation efficiency, time reduction and cost efficiency.

Specifically, samples containing low copy numbers of HIV-1 defined above were used to isolate RNA by the GG method of the present invention described above except that 1 µl each of glycogen at a concentration of 0, 5, 10, 50 or 100 mg/ml was used and the concentration was changed to a final value of 0, 0.1, 0.2, 1 or 2 mg/ml. Isolated RNA was amplified over 30 or 40 cycles of PCR. The primers used for amplification were a gag primer pair of SK 100 and SK 104 and a pol primer pair of P5 and P6. Theoretically, it is expected that 291 bp and 142 bp of PCR products are obtained with the gag and pol primer pairs, respectively.

One µl of a dye was added to 9 µl of the second PCR product and the mixed product was loaded on a 5% polyacrylamide or 2.0% agarose gel and electrophoresed.

Bands were detected by ethidium bromide staining.

The results are shown in Fig. 1. In Fig. 1, panels A. B and C show the results after 30 cycles with the gag

primer pair, 40 cycles with the gag primer pair and 30 cycles with the pol primer pair, respectively. Lanes 1-5 in each panel represent bands at glycogen concentrations of 0 mg/ml, 0.1 mg/ml, 0.2 mg/ml, 1 mg/ml and 2 mg/ml.

5 respectively. Lane 6: positive control (P); lane 7: negative control (N); M: molecular weight marker (Roche).

As shown in Fig. 1, bands were detected at glycogen concentrations of 0.2 mg/ml to 2 mg/ml (lanes 3-5) in each case. Therefore, glycogen can function as a carrier at a 10 concentration of about 0.2 mg/ml or more.

Example 3: Optimal concentration of guanidine thiocyanate used for RNA isolation

The concentration of guanidine thiocyanate added to 70% ethanol during the final ethanol washing step was 15 changed to determine the optimal concentration from the viewpoint of isolation efficiency.

Specifically, samples containing low copy numbers of HIV-1 defined above were used to isolate RNA by the GG method of the present invention described above except that the concentration of guanidine thiocyanate added to 70% ethanol during the final ethanol washing step was changed to 0.0 M, 0.6 M, 0.7 M, 0.8 M, 0.9 M and 1.0 M.

As a result, small amounts of contaminants remained as masses in the final RNA at a guanidine thiocyanate concentration of 0.8 M or less but disappeared at 0.9 M or more.

Example 4: Comparison of RNA isolation efficiency

The efficiency of the GG method of the present

25

invention was compared with those of commercially available known RNA or DNA isolation kits on HIV samples.

Specifically, seven widely used commercial RNA or DNA isolation kits were adopted as controls to perform nucleic 5 acid isolation according to the ordinary protocol of each kit. For each kit, 26 high copy-number samples and 47 low copy-number samples described above were used. Nucleic acids were amplified by PCR with the gag primer pair and/or pol primer pair described above and detected as described above.

The results are shown in Table 5 below.

Table 5

	Isolation	GG method	A	В	С	D
	method					
15	High copy	26/26	10/26	12/26	12/26	10/26
		26/26	10/26	12/26	12/26	10/26
		26/26	11/26	11/26	12/26	9/26
		(100)	(39.7)	(44.9)	(46.2)	(37.2)
20	Low copy	38/47	0/47	0/47	0/47	0/47
		38/47	0/47	0/47	0/47	0/47
		37/47	0/47	0/47	0/47	0/47
		(80.2)	(0)	(0)	(0)	(0)

10

	Isolation	E	F	G
	method			
	High copy	10/26	26/26	26/26
		10/26	26/26	26/26
5		10/26	26/26	26/26
		(38.5)	(100)	(100)
			0	
	rom coba	0/47	33/47	36/47
		0/47	33/47	36/47
10		0/47	32/47	35/47
		(0)	(69.5)	(75.9)

In high copy-number samples, detection was about 35% to about 50% with control kits A to E and 100% with F or G

15 kit. In contrast, the method of the present invention achieved detection of 100%. In low copy-number samples, detection was 0% with control kits A to E and about 70% with F or G kit in contrast to the method of the present invention that achieved detection of 80.2% even in low copy-number samples.

This demonstrates that the present invention shows significantly excellent isolation efficiency as compared with many existing RNA isolation methods.

Example 5: Identification of the sequences of amplified

25 nucleic acids

Amplified samples were sequenced.

Specifically, an automated sequencer ABI-PRISM® 310 (Perkin-Elmer) was used for direct sequencing as

И

recommended by the manufacturer. The target region in this Example is the gag or pol region of HIV-1. The reference sequence was retrieved from the annual report of the Los Alamos National Laboratory. Phylogenetic trees were used for data analysis.

The results on PCR products using the gag primer pair are shown in Fig. 3 and SEQ ID NOS: 10-19. The nucleotide sequences of nucleic acids can be determined in all the samples subjected to RNA isolation by the GG method of the present invention. Sequencing of 10 samples revealed a common sequence of subtype E (Fig. 3). Sequence data show that all the 10 samples tested have a 90% or more homology with the common sequence (SEQ ID NO: 9) in the gag region of HIV-1 (291 bp). In Fig. 3, bases identical to those of the common sequence are shown by "-".

This verifies that each sample subjected to nucleic acid isolation contains HIV-1 RNA and that the presence of HIV-1 was detected by isolation and amplification of the RNA.

20 EFFECT OF THE INVENTION

10

15

We compared known isolation methods using commercial kits with a method of the present invention. The results showed that the method of the present invention was the most excellent isolation method from the viewpoints of detection sensitivity, economy and time reduction.

Conventional methods using CsCl, or a resin were suitable in terms of time reduction, but inferior to the method of the present invention in detection sensitivity.

The method of the present invention is safer than the AGPC method because it uses neither phenol nor chloroform.

Moreover, the necessary period is within 60 minutes, which is shorter than required by conventional methods because any organic solvents such as phenol and chloroform are not used. The method of the present invention can be performed more inexpensively on small amounts (about 50 µl) of samples. All the steps of the method of the present invention can be performed in a 0.5 ml tube with no need to change the tube. Thus, processes from nucleic acid isolation to amplification can be completed in a single step, whereby the risk of contamination of reaction products is decreased.

Without being bound to any theory, the excellent

isolation efficiency of the method of the present invention
is partially attributed to the use of glycogen as a carrier
of nucleic acids. The present invention is particularly
useful for screening small amounts of samples such as blood
products or blood samples from children, for example, and
for following up patients under AIDS therapy, and it is
useful for not only the field of research but also clinical
applications.

CLAIMS

- A nucleic acid isolation kit comprising a reducing agent, a coprecipitant and a protein denaturant characterized in that it contains no protease.
- The nucleic acid isolation kit of Claim 1, which is salt-free.
- The nucleic acid isolation kit of Claim 1 or 2 wherein the nucleic acid is RNA.
- 4. The nucleic acid isolation kit of any one of Claims 1 to 3 wherein the reducing agent is 2-mercaptoethanol or dithiothreitol.
- 5. The nucleic acid isolation kit of any one of Claims 1 to 4 wherein the coprecipitant is glycogen or dextran.
- 6. The nucleic acid isolation kit of any one of Claims 1 to 5 wherein the protein denaturant is guanidine thiocyanate.
- 7. A method for isolating a nucleic acid from a biological sample, comprising:
- i) incubating the biological sample with a reducing agent, a coprecipitant and a protein denaturant to degrade and denature proteins and other contaminants in the biological sample without using a protease, and
- ii) directly performing alcohol precipitation with a lower alcohol.
- 8. The method of Claim 7 wherein incubation takes place at 55°C to 65°C for 5 minutes to 15 minutes in step 1).
- The method of Claim 8 wherein incubation takes place at about 60°C for about 10 minutes in step 1).

- 10. The method of any one of Claims 7 to 9 comprising adding a protein denaturant during alcohol precipitation.
- 11. The method of any one of Claims 7 to 10 wherein the biological component is a body fluid or a blood product.
- 12. The method of any one of Claims 7 to 10 wherein the biological sample has a volume of 30 μ l to 100 μ l.
- 13. The method of any one of Claims 7 to 11 comprising no step of adding a salt.
- 14. The method of any one of Claims 7 to 12 wherein said steps are performed in a single 0.5 ml tube.

ABSTRACT

The present invention provides a nucleic acid
isolation kit comprising a reducing agent, a coprecipitant
and a protein denaturant characterized in that it contains
no protease, and a method for isolating a nucleic acid
using said kit.

Fig.1

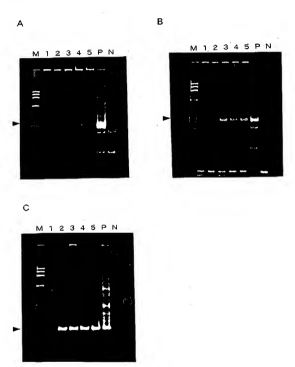


Fig.2

FLOWCHART OF THE GG METHOD

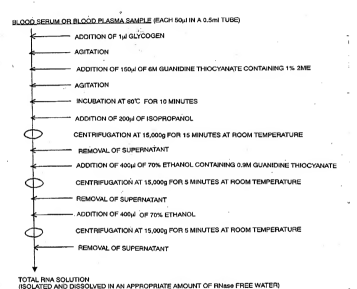


Fig.3

	9111111111	140	\$
	CAGTACALG	-	A Packet
;_		0,1	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
	99 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		G C C C C C C C C C C C C C C C C C C C
;-	00000000000000000000000000000000000000	130	130 130 130 130 130 130
	AG 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0		20079900
+	92	1 10	180 180 180 180 180
	AgGAag		ACCAGE CCACCE
	accatcantg	190	Agardadada Acchagagda agardaaaraha Caacaaraha Tacaacaraha Caacaaraha Tacaacaraha Caacaacaraha Caacaacaacaraha Caacaacaacaraha Caacaacaacaraha Caacaacaacaacaacaacaacaacaacaacaacaacaac
	gtTadagatt	8 <u>-</u>	1 160 1 160
	CEATGCAARY gtraakank accarcand agokageroc Assaryachric	8 -	Some Aggregation Aggrega
	UBICE E		1 1 1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
	COMMON SECURICE E SAMPLE 1 SAMPLE 2 SAMPLE 3 SAMPLE 6 SAMPLE 6 SAMPLE 6 SAMPLE 6 SAMPLE 6 SAMPLE 7 SAMPLE 7 SAMPLE 7 SAMPLE 9		SAMPLE 1 SAMPLE 5 SAMPLE 6 SAMPLE 8 SAMPLE 8 SAMPLE 8 SAMPLE 1 SAMPLE 1 SAMPLE 1 SAMPLE 2 SAMPLE 2 SAMPLE 2 SAMPLE 6 SAMPLE 6 SAMPLE 6 SAMPLE 7 SAMPLE 6 SAMPLE 7 SAMPLE 6 SAMPLE 7 SAMPLE 7 SAMPLE 7 SAMPLE 7 SAMPLE 7 SAMPLE 8 SAMPLE 8 SAMPLE 8 SAMPLE 9 SAM

	•					
					Nbon & Vanderhye	P.C. (12/97)
Case No.			RULE 63 (37 C.F.R. 1.63)			
		DECLAR	ATION AND POWER OF A	TTORNEY		
		IN THE INSTER	FOR PATENT APPLICATION	DEMARK OFFICE		
I am the o	riginal, first and sole inve	enter (it only one name is it	sted below) of all original, it	at and joint intection (ed below next to my name, if plural names are listed be ACID BY USING TH	
KITS F	ication of which (check a	nonlicable box(s)):	-			
is a	ttached hereto					
	s filed on	0.30	as U.S. Application S /JP00/05170		gust 2, 2000	
DCI was	s filed as PCT Internation			on _A	gust 2, 2000	
		pplication) was amended o				
amendme 37 C.F.R. below and priority is Priority Fo Application	ent referred to above. I a 1.58. I hereby claim for I have also identified be claimed or, if no priority preign Application(s): on Number	cknowledge the duty to dis	close information which is n 35 U.S.C. 119/365 of any li for patent or Inventor's certification date of this application:	national to the patenta	the claims, as amended by billity of this application in ac or petent or invertor's certificate before that of the application Day/Month/Year File 5/7/1999	cate listed stion on which
190	633/1999		Japan		31,122,	
i hereby o	laim the benefit under 3 on Number	5 U.S.C. §119(e) of any Un	ited States provisional appli Date/Month/Year Filed	ication(s) listed below		
I hereby o	claim the benefit under 3 atter of each of the clair 12, I acknowledge the du	5 U.S.C. 120/365 of all prices of this application is not	r United States and PCT In disclosed in such prior appl	ternational application ications in the manne	s listed above or below and, r provided by the first paragr ad between the filling date of	Insofar as the aph of 35
U.S.C. 11 applicatio	ns and the national or P	ty to disclose material imor CT international filing date		H. 1.56 WHICH OCCUR		o o pino
applicatio	ns and the national or P	ty to disclose material infor CT international filing date		.H. 1.56 WIEGI OCCU	Ste	tus: patented
application	ns and the national or P JPCT Application(s):	ty to disclose meterial imor CT international filing date		-H. 1.56 WHICH OCCUR	Ste	
application	ns and the national or P	ty to disclose material infor CT international filing date	of this application:	.H. 1.56 WHEN OCCUR	Ste	tus: patented
Prior U.S Application I hereby of be true; a Imprison application 22201-47 address) connecte 30184; R	ns and the national of P s/PCT Application(e): on Serial No. declare that all statemen and further that these at ment, or both, under See nor ary patient issued i rid, telephone number individually and collects de thorewith and with the obert W. Farts, 31352.2 2333, Legorard C. Mine	ts made herein of my own tements were made with thion 1001 of Title 18 of the hereon. And I hereby apper (703) a16-4000 (to whom way my attorneys to prosecreculting patent. Arthur P. Bichard G. Besha, 22770-14.	Day/Month/Year Filed Knowledge are true and three is knowledge are true and three is knowledge are true and three is knowledge are true and true is knowledge and the Linked States Code and the Int NIXON & VANDERHYE all cammunications are to Crawlorg, 25327: Larry S. Lark E. Rubatawa are, 33383: Jeffry H. Nelson widson, 32489. Alen M. Kd	t all statements made to statements and the at such willful faile sit P.C., 1100 North Git be directed), and the ansact all business in tionn_2584(i); Robert 1, iichael J. Keenan, 32*	on Information and belief ar like so made are purishable the Bid. 8° Floor, Julingto et lottlever and the Patent and Trademon's the Patent and Trade	tus: patented g, abandoned e belleved to b by fine or velidity of the n, VA f (of the same office 51; Stanley C. nam. Jr. 29384;
application Prior U.S Application I hereby of the true; a simprison application applicatio	ns and the national of P s/PCT Application(e): on Serial No. declare that all statemen and further that these at ment, or both, under See nor ary patient issued i rid, telephone number individually and collects de thorewith and with the obert W. Farts, 31352.2 2333, Legorard C. Mine	its made herein of my own its made herein of my own its made with the tion 1001 of Title 18 of the hereon. And I hareby apper (703) 816-4000 (to whom reaching patent. Anthur R. Bighand G. Besha, 22770-14 hardt. 29009: Duane M. By Williams 20908: J. Scott D. Williams 20908: J. Williams 20908: J. Williams 20908	Day/Month/Year Filed Knowledge are true and three te knowledge are true and three te knowledge have the true to the knowledge are true and the te knowledge have been to knowledge and the NAON & VANDERHYE all cammunications are to control of the true the application and to to the spice of the true true true the true to the true the true to the true	tall statements made to statements and the at such willful false size D.C., 1100 North Gis be directed), and the ansact all business in liston, 25640; Robert J. 25640; Robert J	on Information and ballet as like as or made are purishtable be Rd., a [®] Floor, Airlingto e following altomyty the Parent and Trademark L. Vandenhye, 27078. Jenne Ce. Byan H. Deviston, 302 Ger, 33149 H. Warner Bur. (Gilfin, 1726), Nober A. k. Deto: April D. Nober A. k.	tus: patented g, abandoned e belleved to a by fine or validity of the n, VA (of the same office of the same of the office of the
Prior U.S Application I hereby of be true; a Imprison application 22201-47 address) connecte 30184; R	ns and the national or P is April and in the national or P is April and in the	te made herein of my own termade herein of my own termade with the tion 1001 of Title 18 of the hereon. And I hereby appe (703) at 8-4000 flo whom any my attempts of prosecution of the hereon herein of the hereon herein of the hereon herein of the herein own the herein of the herein own termade herein own the herein own termade herein	Day/Month/Year Filed Day/Month/Year Filed Menowledge are true and the le incovidedge that without the INCOV A VANDEHIYE the INCOV A VANDEHIYE the INCOV A VANDEHIYE Tark E, Nebaum, 3234E. Tark E, Nebaum, 3234E. Tark E, Nebaum, 3234E. Mill 3/534. Mill 3/534.	t all statements made to statements and the at such willful false statements and the at such willful false statements. P.C., 1100 North Gil be directed), and the ansast all business in bloom, 258-07; Robert Libral J. Keenera, 32, 30581; John R. Tussigen, 36178; William J. Von Libral J. Keenera, 32, 30581; John R. Tussigen, 36178; William J. Von Libral J. Keenera, 32, 30581; John R. Tussigen, 36178; William J. Von Libral J. Von Libra	Site pendin on Information and belief are ible so mede are punishable sements may jeopardize the Rd., 6º Hoor, All Ingrio e following attorneys therein the Peters and Types and the Types and Ty	tus: patented g, abandoned e belleved to by fine or validity of the n, VA (of the same Office 17. Hosmer, 51; Stanley C, nam. Jr. 29306; tolan, 29834;
Prior U.S Application I hereby of be true; a Imprison application 22201-47 address) connecte 30184; R	ns and the national or P London Serial No. doclars that all statomer and further that these as a ment, or both, under Soc on or any patent issued to nor any patent issued to hother W. Faria, 19352: 12 27395, 1eonard C. Mitt. E Byrne, 322055. Many J Lord, 36655, Jannes D. Br Invoentor's Signature: Inventor's Signature: Inventor's Signature:	is made herein of my own terments were made which the herein. And I hereby spotion 1001 of 116 is 8 of the herein. And I hereby spotion 1001 of 116 is 8 of the herein. And I hereby spotion 1001 of 116 is 8 of the herein. And I hereby spotion 1001 of 1001	month application: Day/Month/Year Filed knowledge are mue and the te knowledge that willful risk in the Month of the te knowledge that willful risk int NIKON & VANDERHYE unted Stites Code and the int NIKON & VANDERHYE and communications are to not this application and to the tage is the tage of tage of the tage of tage of the	t all statements made to statements and the statements and the statements and the statements and the statements. P.C., 1100 North Git be directed, and the statement all submisses in soon, 256-06, 760-061. Deads it, with RTLBS and 361-761. William Statement and Statement State	Site pendin on Information and belief are ible so mede are punishable sements may jeopardize the Rd., 6º Hoor, All Ingrio e following attorneys therein the Peters and Types and the Types and Ty	tus: patented g, abandoned e belleved to a by fine or validity of the n, VA (of the same office of the same of the office of the
Prior U.S Application I hereby of be true; a Imprison application 22201-47 address) connecte 30184; R	ns and the national or P "IPCT Application(e): on Serial No. declare that all statement for the threes etc. and rurber that three etc. and rurber that the	ts made herein of my own terminations him gase with a herein of my own termination of the second of Title 1s of the hereon. And I hareby spor (703) s15-4000 (fic whom why my attempts of prose-resulting patent. Arthur R. H. By. Wilson, 32955, J. Scott D. And J. G. Wilson, 32955, J. Scott D. And J. G. Wilson, 32955, J. Scott D. And J. G. Wilson, 32955, J. Scott D. Maniko (J. Wilson, 32955, J. Scott D. Wilso	Day/Month/Year Filed for which application: Day/Month/Year Filed for which application and to the should be application and the should be application and the should be application and to the should be application and the should be applied to t	t all statements modes o statements and the ostatements and the P.C., 1100 North Grant Statements of the Grant Statement of the Grant Sta	on information and belief as like so made are purishable to like so made are purishable to like so the Ad. 8° Floor, All inglot o following attorneys thereon to following attorneys thereon 6°, Bryan H. Deviston, 302 Gro., 35149; H. Warner Burn. Garitin, 31260; Robert A. N. Deto: APT. 1. Deto: APT. 1. Liapanes. (citi. 1.)	tus: patented g, abandoned e belleved to by fine or validity of the n, VA (of the same Office 17. Hosmer, 51; Stanley C, nam. Jr. 29306; tolan, 29834;
Prior U.S Application I hereby of be true; a Imprison application 22201-47 address) connecte 30184; R	ans and the national or P LOTCI Application(e): on Serial No. declare that all statement and further that these at a ment, or both, under Soc on 1-1, elephone number of 1-1, elephone number of thorewith and with the control of the series	is made herein of my own terments were made which the herein. And I hereby spotion 1001 of 116 is 8 of the herein. And I hereby spotion 1001 of 116 is 8 of the herein. And I hereby spotion 1001 of 116 is 8 of the herein. And I hereby spotion 1001 of 1001	Day/Month/Year Filed for only of this application: Day/Month/Year Filed for only of the provided per true and the se innovided per true and the application and the una this application and the una think application and the unapplication and unapplication a	s all statements made o statements and the ostatements and the P.C., 1100 North Gill and the ansatz all business in be directed, Robert (ichael J. Keenan, 32, 39641: John R.T. Lasigen, 36175, Vijikam J. VOSH HARA (1689)	Site pendin on Information and belief are bids so made are punishable sements may jogardize the Rd., 6° Hoor, All Ingto e following attorneys thereon the Pattern and Trademark of Bysan H. Davidson, 30°c, 331-8, H. waren Burr. G. Giffin, 31:260; Robert A. b. Dete: April 1. L. April	tus: patented g, abandoned e belleved to by fine or validity of the n, VA (of the same Office 17. Hosmer, 51; Stanley C, nam. Jr. 29306; tolan, 29834;
Prior U.S Application I hereby of be true; a Imprison application 22201-47 address) connecte 30184; R	ns and the national or P "IPCT Application(e): on Serial No. declare that all statement for the threes etc. and rurber that three etc. and rurber that the	ts made herein of my own terminations him gase with a herein of my own termination of the second of Title 1s of the hereon. And I hareby spor (703) s15-4000 (fic whom why my attempts of prose-resulting patent. Arthur R. H. By. Wilson, 32955, J. Scott D. And J. G. Wilson, 32955, J. Scott D. And J. G. Wilson, 32955, J. Scott D. And J. G. Wilson, 32955, J. Scott D. Maniko (J. Wilson, 32955, J. Scott D. Wilso	Day/Month/Year Filed for which application: Day/Month/Year Filed for which application and to the increasing a second and application and to the care to the second application and to the care of th	t all statements modes o statements and the ostatements and the P.C., 1100 North Grant Statements of the Grant Statement of the Grant Sta	Site pendin on Information and belief ar Bibe so made are punishable temoris may jeopardize the Rid, \$\frac{2}{3}\text{ Fig. 3}\text{ Fig. 3}\	tus: patented g, abandoned e belleved to a by fine or validity of the n, VA (of the same Office 17, Nosmer 5, Stantey C, stann, Jr. 29306; Stantey C, stann, Jr. 29306; 2002
Prior U.S Application I hereby of be true; a Imprison application 22201-47 address) connecte 30184; R	ns and the national or # "PCT Application(e): on Serial No. declare that all statement and further that these as a ment, or both, under Sec ment, or both, and collect discovers the property of t	ts made herein of my own tempers were made with a tempers were made with a tempers were made with a tempers, and therein, and therein, and therein and tempers are made with a wind and the tempers and the tempers and tempers are the tempers and tempers and tempers are tempers and tempers and tempers are tempers are tempers and tempers are tempers and tempers are tempers are tempers and tempers are tempers and tempers are tempers and tempers are te	Day/Month/Year Filed for which application: Day/Month/Year Filed for which application and to the should be application and the should be application and the should be application and to the should be application and the should be applied to t	t all statements made to statements and the to statements and the to such willful factor per per per per per per per per per pe	Site pendin con information and belief are to be so made any purishtable semants may popardize the ERI, d.** Floor, Allingto e following attorneys thereon the Petern and Trademark (Vandemye, 27075, Stemother, 2	tus: patented g, abandoned e believed to be by fine or velicity of the n, VA (of the same Office 71, Hosmer, 51; Stanley C, 13, 13, 13, 13, 13, 13, 13, 13, 13, 13
application Prior U.S Application I hereby a be true; a imprison application application 22201-47 address) connected 30184; B Spooner, Thomas B. J. Sad	ans and the national or P LapTCT Application(e): on Serial No. doclare that all statement and further that these at a ment, or both, under Sec on or any patent issued 1-d, steephene number instance and the second or any patent issued to be the second or any patent issued to the second or any patent is the second or any pate	ts made herein of my own terminations here of the herein of my own termination from the fine 18 of the hereon. And hereby apper (703) at 8-4000 flow whom any my attempts of prosecution for the first of the first own termination from the first own termination for the first own termination from the first own termination for the first own termination from the first own tex	Day/Month/Year Filed knowledge are true and the elementedge that willful falls the second and the month of the second and the month of the second and the min NKON & VANDEHIYE Lang S. It and the second and the min NKON & VANDEHIYE Conserved, 253251. Juny S. N. Tark E. Nedbaum, 353251. Juny S. N. Tark E. Nedbaum, 353251. Juny S. N. Martina L. Nedbaum, 353251. Juny S. N. Martina L. Nedbaum, 353251. Juny S. N. Martina L. N. Toky S. M. Z. L.	s all statements made to statements and to statement	Site pendin on Information and belief are like so made are punishable sements may jeopardize the Rid., 8° Hoor, All Ingrio to following attorneys therein the Petiden are 2007.8 Jenne Ces. Bryan H. Devidson, 2007. 33149. H. Warron Burr. G. Griffin, 31260; Robert A. k. Dete: April 1. J. Japanes. Japan. Date: April 1. J. Japanes. April 1. J. Japanes. Jap	tus: patented g, abandoned e bolleved to a by fine or validity of the n, VA (of the same office of the same
application Prior U.S Application I hereby a be true; a imprison application application 22201-47 address) connected 30184; B Spooner, Thomas B. J. Sad	ns and the national or P "PCT Application(e): on Serial No. declare that all statement of both under Sec ment, or both, under Sec production of the second	ts made herein of my own terments were made with the terment to the termen	of this application: Day/Month/Year Filed knowledge are true and their less from the first willful risk less from the first willful risk less from the first willful risk less from the first NIKON & VANDERHYE and communications are to no this application and to another special communication and to not the application and to another first E. Rubatum 32348. Mark E. R	t all statements made to statements and the to statements and the to such willful factor per per per per per per per per per pe	Site pendin on information and belief are pendin to be a made are punishable senants may be pendin the Rd. of "Hoor All inglot of following attorneys thereon the Peters and Trademark Vandering, 27076, Jenne Vandering, 2707	tus: patented g, abandoned e believed to be by fine or velicity of the n, VA (of the same Office 71, Hosmer, 51; Stanley C, 13, 13, 13, 13, 13, 13, 13, 13, 13, 13
application Prior U.S Application I hereby a be true; a imprison application application 22201-47 address) connected 30184; B Spooner, Thomas B. J. Sad	ns and the national or # "PCT Application(e): on Serial No. declare that all statement and further that these as a ment, or both, under Sec ment, or both, and collect discovers the property of t	ts made herein of my own terminations here of the herein of my own termination from the fine 18 of the hereon. And hereby apper (703) at 8-4000 flow whom any my attempts of prosecution for the first of the first own termination from the first own termination for the first own termination from the first own termination for the first own termination from the first own tex	Day/Month/Year Filed knowledge are the semination of the september of the semination	t all statements made to statements and the total country of the statements and the total country of the statements and the total country of the statement of t	Site pendin on Information and belief are like so made are punishable sements may jeopardize the Rid., 8° Hoor, All Ingrio to following attorneys therein the Petiden are 2007.8 Jenne Ces. Bryan H. Devidson, 2007. 33149. H. Warron Burr. G. Griffin, 31260; Robert A. k. Dete: April 1. J. Japanes. Japan. Date: April 1. J. Japanes. April 1. J. Japanes. Jap	tus: patented g, abandoned e bolleved to a by fine or validity of the n, VA (of the same office of the same

FOR ADDITIONAL INVENTORS, check box 🚺 and attach sheet with same information and signature and date for each.

.6309-3. Komuro, Ina-machi, Kitaadachi-gun, Saitama

Residence: (city)
Post Office Address:
(Zip Code)

Japanese

Case No.

Post Office Address: (Zip Code) . Nixon & Vanderhye P.C. (12/97)

RULE 63 (37 C.F.R. 1.63) DECLARATION AND POWER OF ATTORNEY FOR PATENT APPLICATION IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

1 1, 2002 inventor's Signature: MANABE Sachiko Inventor: (citizenshlp) (first) (state/country) Residence: (city) Tokyo (state/country) Iapan 2-1-12-803, Kitaueno, Taito-ku, Tokyo 110-0014 Japan Post Office Address: · (Zip Code) Date: Inventora Signature: Inventor: (citizenship) (state/country) Residence: (city) Post Office Address: (Zip Code) Date: Inventor's Signature: Inventor: (citizenship) (first) (iast) (state/country) Residence: (city) Post Office Address: (Zip Code) Date: 7. Inventor's Signature: Inventor: (first) (last) (state/country) Residence: (city) Post Office Address: (Zip Code) Date: Inventor's Signature: inventor. (citizenship) (first) (fast) (state/country) Residence: (city)
Post Office Address:
(Zlp Code) Inventor's Signature: 9. Inventor (tast) (cltizenship) (first) (state/country) Residence: (city) Post Office Address: (Zip Code) Date: Inventor's Signature: 10. Inventor (citizenship) (state/country) Residence: (city) Post Office Address: (Zip Code) Date: inventor's Signature: 11. inventor: (last) (first) (state/country) Residence: (city) Post Office Address: (Zip Code) Date: Inventor's Signature: 12. Inventor: (last) (citizenship) (state/country) Residence: (city)

SEQUENCE LISTING

<1	10>	Oriental Yeast Co., Ltd.	
		NAITOU Toshikuni	•
		National Institute of Infectious Diseases	
		TAKEDA Yoshifumi	
<1	20>	Kits for extracting nucleic acid and method of extracting	•
nu	clei	c acid by using the kits	
<1	30>	YCT513	
<1	60>	19	
<2	10>	1 ,	
<2	11>	23	
<2	12>	DNA	
<2	13>	Artificial Sequence	
<4	<00	1	
aa	ggat	agag gtaaaagaca cca 23	
<2	10>	2	
<2	11>	23	
<2	12>	DNA	
<2	13>	Artificial Sequence	
<4	<00	2	
t a	igctg	getgg tecaatgett tta 23	
<2	210>	3	
<2	211>	19	
<2	212>	DNA	
0	213>	Artificial Sequence	

(100)	
atcaagcagc catgcaaat 19	
⟨210⟩ 4	
<211> 22	
<212> DNA	
<213> Artificial Sequence	
<400> 4	
cilliggtee tigicitate te 22	
<210> 5	
<211> 23	
<212> DNA	
<213> Artificial Sequence	
<400> 5	
agiggatata tagaagcaga agt 23	
<210> 6	
<211> 27	
<212> DNA	
<213> Artificial Sequence	
<400> 6	
eccecatice eccettitet tilaaaa	27
<210> 7	
<211> 18	
<212> DNA	
<213> Artificial Sequence	
<400> 7	

attagcagga agatggcc	18		
(2.42)			
⟨210⟩ 8			
<211> 18			
<212> DNA			
<213> Artificial Sequ	ence		
<400> 8			
tactccttga cttlgggg	18		
<210> 9			
<211> 220			
<212> DNA		Y	
<213> HIV-1			
<400> 9			
ctatgcaaat gttaaaagal	accalcaalg aggaagelge	agaalgggac	50
agggtacatc caglacaigc	agggeetatt ceaceaggee	agatgagaga	100
accaagggga agtgacatag	caggaactac tagtaccctt	caagaacaaa	150
taggatggat gacaagcaat	ccacctatcc cagtgggaga	catctataaa	200
agatggataa tootgggatt			220
<210> 10			
<211> 220			
. <212> DNA			
<213> HIV-1			
<400> 10			
ccatgcaaat gttaaaagaa	accatcaatg aggaagctgo	agaatgggat	50
agggtacacc cagtacaigc	agggcciail ccaccaggc	aaalgaggga	100
accaagggga agtgacatag	caggaactac tagtaacct	caagaacaaa	150
taggatggat gacaagcaat	ccacctatcc cagtgggag	a catctataaa	200

aggtggata	a teetgggatt				220
	5.				
<210> 11					
<211> 22	0				
<212> DN	A				
<213> HI	V-1				
<400> 11					
ccatgcaaa	t gitaaaagaa	accalcaatg	aggaagctgc	agaatgggat	50
agggtacac	c caglacatge	agggcctatt	ccaccaggcc	agalgaggga	100
accaagggg	a agigacalag	cagggactac	tagtaacctt	caagaacaga	150
taggatgga	i gacaaacaat	ccacctatcc	cagigggaga	catclalaaa	200
aggtggata	a teetgggatt				220
<210> 12					
<211> 22	0				
<212> DN	IA				
<213> HI	V-1				
<400> 12	!				
ccatgcaaa	it gilaaaagaa	accatcaatg	aggaagctgc	agaatgggat	50
agggtacad	c cagtacatge	agggcctatt	ccaccaggcc	agalgaggga	100
accaaggg	ga agtgacatag	caggaactac	tagtaccett	caagaacaaa	150
taggatgg	at gacaaacaat	ccacctatcc	cagtgggaga	catcialaaa	200
agatggat	aa toolgggali				220
<210> 1	3				
<211> 2	20				
<212> D	NA				
<213> H	I V-1				
(400) I	3				

ccaigcaaai	gilaaaagaa	accatcaatg	aggaagctgc	agaatgggat	50
agggtacacc	cagtacatgo	agggcctatt	ccaccaggcc	agatgaggga	100
accaagggga	agigacatag	caggaactac	tagtaccctt	caagagcaaa	150
taggatggat	gacaaacaat	ccacctatcc	cagtgggaga	catctataaa	200
aggtggataa	tcctgggatt				220
<210> 14					
<211> 220					
<212> DNA					
<213> HIV-	-1				
<400> 14					
ccatgcaaat	gttaaaagaa	accatcaatg	aggaagctgc	agaatgggat	50
aggitacacc	caatacatgc	agggcctatt	ccaccaggcc	agatgaggga	100
accaagggga	agtgacatag	caggaaccac	tagtaccctt	caagaacaaa	150
taggatggat	gacaaacaat	ccacctatcc	cagtgggaga	calclalaaa	200
aggtggataa	tectgggatt				220
<210> 15					
<211> 220					
<212> DNA					
<213> HIV-	-1				
<400> 15					
ccaigcaaat	gttaaaagaa	accatcaatg	aggaagetge	agaatgggal	50
agggtacacc	cagtacatgo	agggcctatt	ccaccaggco	agalgaggga	100
accaagggga	agtgacatag	caggaactac	tagtaccctt	caagaacaaa	150
taggatggat	gacaaacaat	ccacctatco	cagtgggaga	catctataaa	200
aggtggataa	tcctgggatt				220

<210> 16

<211>	220					
<212>	DNA					
<213>	HIV-	-1				
<400>	16					
ccatgo	aaat	gttaaaagag	accatcaatg	aggaagctgc	agaatgggat	50
agggta	cacc	cagtacatgc	agggcctatt	ccaccaggcc	agalgaggga	100
accaag	ggga	agigatalag	caggaaclac	tagtacccit	caagaacaaa	150
taggat	gga t	gacaagcaat	ccacctatcc	cagtgggaga	catctataaa	200
aggtgg	ataa	tcctgggatt				220
<210>	17					
<211>	220	÷				
<212>	DNA					
<213>	HIV-	-1				
<400>	17			-		
ccalgo	aaat	gilaaaagaa	accatcaatg	aggaagctgc	agaalgggal	50
agggta	cacc	caglacalgo	agggcclali	ccaccaggac	agalgaggga	100
accaag	ggga	agigacaiag	caggaaclac	lagiacccii	caagaacaaa	150
taggat	ggat	gacaagcaal	ccatctatcc	cagigggaga	aaiciaiaaa	200
aggtgg	ataa	tcctgggatt				220
<210>	18					
<211>	220					
<212>						
<213>	HIV	-1				
<400>	18					
ccatgo	aaat	gttaaaagaa	accatcaatg	aggaagctgc	agaatgggat	50
agggta	LCACC	cagtacatgo	agggcctatt	ccaccaggcc	agatgaggga	100
200220	10000	agtgacatag	caggaactac	tagtaccett	caagaacaga	150

taggatggal	gacaaacaat	ccacctaicc	cagigggaga	catctataaa	200	
aggtggataa	tcctgggatt				220	
<210> 19						
<211> 220						
<212> DNA						
<213> HIV	-1					
<400> 19						
ccatgcaaat	gttaaaagac	accatcaatg	aggaagctgc	agaatgggat	50	
agggtacacc	cagtacatge	agggcctatt	ccaccaggcc	agatgaggga	100	
accaagggga	agigacalag	caggaactac	tagtaccctt	caggaacaaa	150	
taggatggat	gacaaacaal	ccacctatcc	cagigggaga	catctataaa	200	
aggtggataa	tcctgggatt				220	

United States Patent & Trademark Office Office of Initial Patent Examination -- Scanning Division



pplication deficien	cies found during	scanning:	
☐ Page(s)	of		were not presen
for scanning.		(Document title)	
		•	
□ Page(s)	of		were not
present for scanning.		(Document title)	

p Scanned copy is best available. Downing tig is dosk